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Remarks:

This application was filed on 15/10/2003 as a divisional application to the application mentioned under INID code 62.

- (54) Novel formulation containing paroxetine
- (57) Controlled release and delayed release formulations which are adapted or intended for oral administration and which contain paroxetine hydrochloride, such as a formulation which is a system for the controlled and delayed release of paroxetine hydrochloride, having (a) a deposit-core comprising 28.61 mg paroxetine hydrochioride, 18.7 mg methocel K4M, 79.14 mg lactose monohydrate, 2.50 mg polyvinylpyrrolidone, 1.25 mg magnesium stearate, and 0.50 mg Syloid 244.

and (b) a support-platform applied to said deposit-core comprising 15.04 mg Compritol 888, 30.50 mg lactose monohydrate, 4.00 mg polyvinylpyrrolidone, 0.80 mg magnesium stearate, 29.32 mg methocel E5, 0.32 mg Syloid 244, and 0.02 mg iron oxide; and (c) an enteric coating comprising 13.27 mg Eudragit, 3.31 mg talc, and 1.33 mg triethyl citrate, may be used to treat a variety of disorders including depression.

Description

- [0001] The present invention relates to a novel formulation containing paroxetine or a pharmaceutically acceptable salt thereof, and to its use in the treatment and/or prophylaxis of certain disorders.
- [0002] US Patent No 4,007,196 describes inter alia a compound which is commonly known as paroxetine. This compound is a Selective Serotonin Regulake Inhibitor (SSRI) and is currently marketed world-wide for the treatment and/or proph/axis of depression.
- [0003] The current formulation which is the only marketed formulation of paroxetine hydrochloride is a swallow tablet. [0004] It has now been surprisingly found that controlled release and delayed release formulations containing paroxetine diver isso to an unexpected reduction in the side effects associated with swallow tablets.
- [0005] Accordingly, the present invention provides a controlled release or delayed release formulation containing paroxetine or a pharmaceutically acceptable salt thereof.
- [0006] A further aspect of the invention provides a controlled release or delayed release formulation containing an SSRI. Examples of SSRIs other than paroxetine include fluovetine (US Patent No. 4,3874,081), fluvoxamine (US Patent No. 4,085,225), and sertraine (US Patent No. 4,085,225), and sertraine (US Patent No. 4,595,618).
- [0007] By controlled release is meant any formulation technique wherein release of the active substance from the dosage from is modified to occur at a slower rater than that from an immediate release product, such as a conventional
- swallow tablet or capsule.

 [0008] By delayed release is meent any formulation technique wherein release of the active substance from the
 dosage form is modified to occur at a later time than that from a conventional immediate release product. The subse-
- quent release of active substance from a delayed release formulation may also be controlled as defined above. [0009] Examples of controlled release formulations which are suitable for incorporating paroxetine and other SSRIs are described in:
- 26 Sustained Release Medications, Chemical Technology Review No. 177. Ed. J.C. Johnson. Noyes Data Corporation 1980
 - Controlled Drug Delivery, Fundamentals and Applications, 2nd Edition. Eds. J.R. Robinson, V.H.L. Lee. Mercel Dekkes Inc. New York 1987.
- 30 [0010] Exemples of delayed release formulations which are suitable for incorporating paroxetine and other SSRIs are described in:
 - [0011] Remington's Pharmaceutical Sciences 16th Edition, Mack Publishing Company 1980, Ed. A. Osol.
- [0012] Such controlled release formulations are preferably formulated in a manner such that release of active substance such as paroxienis in effected precionimating during the passage through the stomach and the small intestline, and dalayed release formulations are preferably formulated such that release of active substance such as paroxeline is avoided in the stomach and is effected predominantly during passage through the small intestline
- [0013] Said formulations are preferably formulated such that the release of the active substance is predominantly 1½ to 3 hours post ingestion.
- [0014] The small intestine is suitably the duodenum, the ileum or the jejunem.
- 40 [0015] Petients who benefit most from the formulations of the present invention are those who are known to suffer from nausea upon oral administration using swallow tablets.
 - [0016] Preferred formulations are ultimately enteric coated tablets or caplets, wax or polymer coated tablets or caplets or time-release matrices, or combinations thereof.
 - [0017] Particularly preferred formulations are described in US Patent No. 5,102,666.
 - 6 [0018] Thus, a particular aspect of the invention provides a polymeric controlled release composition comprising a reaction complex formed by the interaction of (f) a calcium polycarbophil component which is a water-swellable, but water insolute, fitnous cross-inked cateoxy-functional polymer, said polymer containing (a) a plurality of repeating units of which at least about 80% contain at least one carboxyl functionality, and (b) about 0.05 to about 1.5% cross-linkina aeant substantially free from polyalkenyl ovighers, said preventages being based upon the weights of unpoly-
- inking agent aubstantially free from polyatenyl polyateners, sain percentiages being passe upon un evergins or unpoymented repealing unit and cross-linking agent, respectively, with (2) water, in the presence of an active agent selected from the group consisting of SSRIs such as paroxetine. The amount of calcium polycarbophil present is from about 0.1 to about 95% by weight, for example about 19%. The amount of active agent present is from about 0.0001 to about 65% by weight, for example between about 5 and 20%. The amount of water present is from about 5 to about 200% by weight, for example between about 5 and 05%. The interaction is carried out at a pl of between about 3 and about
- 55 10, for example about 6 to 7. The calcium polycarbophil is originally present in the form of a calcium salt containing from about 5 to about 25% calcium.
 - [0019] Further particularly preferred formulations are described in US Patent No. 5,422,123.
 - [0020] Thus, a further particular aspect of the invention provides a system for the controlled release of an active

substance which is an SSRI such as parovetine, comprising (a) a deposit-core comprising an effective amount of the active substance and having defined geometric form, and (b) a support-platform applied to said deposit-core, wherein said deposit-core contains at least the active substance, and at least one member selected from the group consisting of (1) a polymeric material which swells on contact with water or a queueus liquids and a geliable polymeric material to said geliable polymeric material wherein the ratio of the said swellable polymeric material is asid geliable polymeric material having both swelling and gelliap properties, and wherein the support platform is an elastic support, applied to said deposit-core so that it partially covers the surface of the deposit-core and follows changes due to hydration of the deposit-core and is slowly soluble and/or slowly geliable in aqueous false. The support-platform may comprise polymers such as hydroxypropylmethylealfulose, plasticizers such as a glycarde, binders such as polymylpyrrollome, hydrophilic agens such as such as such as such as such as a such as a disposit of the support-platform, for example about 15 CoVB. Binder such 40 to 50 Ks. Binder s

15 [0021] Paroxetine used in the present invention is suitably in the form of the free base or a pharmaceutically acceptable salt thereof. Preferably, paroxetine is suitably in the form of the hydrochloride hemitydrate.

[0022] Paroxetine hydrochloride hemihydrate may be prepared according to the procedures generally outlined in US Patent 4,721,723.

[0023] Paroxetine in the form of a controlled release or delayed release formulation can be used to treat and prevent the following disorders:

Alcoholism Anxiety

Depression Obsessive Compulsive Disorder

Panic Disorder Chronic Pain

Obesity Senile Dementia Migraine

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Bulimia Anorexia Social Phobia

Pre-Menstrual Syndrome (PMS)
Adolescent Depression

Trichotiliomania Dysthymia Substance Abuse

40 [0024] These disorders are herein after referred to as "the disorders".

[0025] The present invention provides a method of treating and/or preventing the disorders by administering an effective and/or a prophylactic amount of a controlled release or delayed release formulation containing paroxetine or a pharmaceutically acceptable sail thereof, to a sufferer in need thereof.

[0025] The present invention further provides the use of a controlled release or delayed release formulation contains in paroxetine or a pharmaceutically acceptable salt thereof in the manufacture of a medicament, for treating and/or preventing the disorders.

[0027] The present invention also provides a pharmaceutical composition for use in the treatment and/or prevention of the discorders which comprises a controlled release or delayed release formulation containing paroxetine or a pharmaceutically acceptable sail thereof.

[50 [0028] The following examples illustrate the present invention.

Example 1 (Hydrophilic Matrix)

[00291

Intragranular	% w/w
Paroxetine Hydrochloride	11.45

(continued)

Intragranular	% w/w
Methocel E5	1.25
Lactose	12.3
Extragranular	
Methocel K100LV	30.0
Lactose	44.0
Magnesium Stearate	1.0
TOTAL	100.0

Example 2 (Hydrophilic Matrix)

[0030]

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Intragranular	% w/w
Paroxetine Hydrochloride	11.45
Methocel E5	1.25
Lactose	12.3
Extragranular	
Methocel K100LV	27.5
Methocel K4M	7.5
Lactose	39.0
Magnesium Stearate	1.0
TOTAL	100.0

Example 3 (pH Sensitive Coat on Immediate Release Core)

[0031]

Tablet Core	%w/w
Paroxetine Hydrochloride	11.45
Lactose	64.05
Microcrystalline Cellulose	20.0
Sodium Starch Glycollate	4.0
Magnesium Stearate	0.5
TOTAL	100.0

Tablet Coating (apply approximately 6-10% of tablet core weight)	%w/w	
Hydroxypropylmethylcellulose Phthalate		
Triacetin	10.0	

50 Example 4 (pH Sensitive Coat on Immediate Release Core)

[0032] Tablet Core as in Example 3

Tablet Coating (apply approximately 6-10% of tablet core weight)		
Cellulose Acetate Phthalate	90.0	
Diethyl Phthalate	10.0	

Example 5 (Controlled Release Coating on Immediate Release Core)

[0033] Tablet Core as in Example 3

Tablet Coating (apply approximately 5-12% of tablet core weight)	
Eudragit RS 100	86.0
Dibutyl Phthalate	10.0
Talc	4.0
FD&C Yellow No. 6	0.01

Example 6 (pH Sensitive Coat on Controlled Release Core.)

[0034] Tablet Core as in Example 3 [0035] Tablet Coating as in Example 3

Example 7 (Encapsulated Controlled Release Coated Beads)

[0036]

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Pellet	%w/w (approx)
Non Pareil Seed	30
Paroxetine Hydrochloride	40
Gelatin	8
Lactose	20
Talc	2

Coating	%w/w
Glycerylmonostearate	36.6
Glyceryldistearate	53.4
White Wax	10.0

Example 8 (Controlled release bilayer tablet)

[0037]

Active Layer		
Component	mg/tablet	Function
Paroxetine Hydrochloride	22.89*	Active
Methocel K4M	15.00	Hydrogel polymer
Lactose monohydrate	62.0	Hydrophilic agent
Polyvinylpyrrolidone	3.0	Binder
Magnesium stearate	1.0	Hydrophobic agent
Syloid 244	1.0	Hydrophilic agent
Support platform		
Component	mg/tablet	Function
Compritol 888	15.04	Plasticizer
Lactose monohydrate	29.32	Hydrophilic agent
Polyvinylpyrrolidone	4.0	Binder

*Equivalent to 20mg paroxetine as free base.

(continued)

Support platform		
Component	mg/tablet	Function
Magnesium stearate	1.52	Hydrophobic agent
Methocel E5	29.32	Hydrogel polymer
Iron oxide	0.08	Colourant
Total tablet weight	184.89mg	

[0038] The powder blend for each layer was wet granulated in a high shear mixer/granulator and dried in a fluid bed drier. The bilayer tablets were compressed on a Manesty triple layer press.

Example 9 (Enteric coated calcium polycarbophil formulation)

[0039]

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Core			
Component	mg/ tablet	Function	
Paroxetine Hydrochloride	22.89*	Active	
Calcium polycarbophil	20.00	Matrix	
Lactose anhydrous	146.11	Hydrophilic agent/diluent	
Polyvinylpyrrolidone	10.0	Binder	
Magnesium stearate	1.0	Hydrophobic agent/lubricant	
Water**	0.024	Granulating liquid	
Enteric coat			
Component	mg/ tablet	Function	
Eudragit	22.19	Polymer	
Talc	1.53	Lubricant	
Triethyl citrate	1.00	Plasticizer	
Water**	24.6	Diluent	
Film coat			
Opadry pink	10.5	Film coat	
Water**	94.5	Diluent	
Polish coat			
Opadry clear	0.750		
Water**	29.3	Diluent	

[&]quot;Equivalent to 20mg paroxetine as free base.

[0040] The core constituents were wet granulated in a high shear mixer/granulator, and dried in a fluid bed drier. The magnesium stearate was then added and the inixture processed in a low shear mixer. The mix was then compressed on a B type roary tablet press. Coaling was carried out using an Accele acta.

[&]quot;Removed during processing.

Example 10 (Controlled release bilayer tablet)

[0041]

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Active Layer mg/tablet Component Function Paroxetine Hydrochloride 22.89* Active Methocel K4M 20.00 Hydrogel polymer Lactose monohydrate 60.0 Hydrophilic agent Polyvinylpyrrolidone 5.0 Binder Hydrophobic agent Magnesium stearate 1.0 1.0 Hydrophilic agent Syloid 244 Support platform

Component	mg/tablet	Function
Compritol 888	14.72	Plasticizer
Lactose monohydrate	30.60	Hydrophilic agent
Polyvinylpyrrolidone	2.80	Binder
Magnesium stearate	0.80	Hydrophobic agent
Methocel E5	30.60	Hydrogel polymer
Syloid 244	0.40	Hydrophilic agent
Iron oxide	0.08	Colourant
Total tablet weight	100 00ma	1

*Equivalent to 20mg paroxetine as free base.

[0042] The process was as described in Example 8.

Example 11 (Controlled release bilayer tablet)

[0043]

ctive Layer		
Component	mg/tablet	Function
Paroxetine Hydrochloride	22.89*	Active
Methocel K4M	15.00	Hydrogel polymer
Lactose monohydrate	63.31	Hydrophilic agent
Polyvinylpyrrolidone	2.0	Binder
Magnesium stearate	1.0	Hydrophobic agent
Syloid 244	0.40	Hydrophilic agent
Support platform - as in Example 10.		
Total tablet weight	184.60mg	

^{*}Equivalent to 20mg paroxofine as free base.

^[0044] The process was as described in Example 8.

Example 12 (Enteric coated controlled release bilayer tablet)

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Active Layer		
Component	mg/tablet	Function
Paroxetine Hydrochloride	28.61*	Active
Methocel K4M	18.75	Hydrogel polymer
Lactose monohydrate	79.14	Hydrophilic agent
Polyvinylpyrrolidone	2.50	Binder
Magnesium stearate	1.25	Hydrophobic agent
Syloid 244	0.50	Hydrophilic agent
Support platform		
Component	mg/tablet	Function
Compritol 888	15.04	Plasticizer
Lactose monohydrate	30.50	Hydrophilic agent
Polyvinylpyrrolidone	4.00	Binder
Magnesium stearate	0.80	Hydrophobic agent
Methocel E5	29.32	Hydrogel polymer
Syloid 244	0.32	Hydrophilic agent
Iron oxide	0.02	Colourant
Enteric coating		
Component	mg/tablet	Function
Eudragit	13.27	Polymer
Talc	3.31	Lubricant
Triethyl citrate	1.33	Plasticizer
Water**	36.25	Dituent
Total tablet weight	228.66mg	

^{*}Equivalent to 25mg paroxetine as free base. "*Removed during processing.

[0046] The process was as described in Example 9.

Example 13

GI tolerance study

Treatment:

[0047] The design of the study is outlined below

Subjects: Normal healthy volunteers

Design: Parallel group, placebo controlled, double blind

(a) Placebo, (b) Immediate release paroxetine, (c) Example 8 formulation, (d) Example 8 for-

mulation with enteric coating.

30 mg once daily for 3 days

Dosage: Number of subjects: 452 evaluable (488 randomised, 485 evaluable)

[0048] The study was conducted to compare the incidence, severity and duration of nausea and vomitting, and diarrhoea (theoretically if the controlled release formulations slow down absorption of paroxetine then, as paroxetine is known to be prokinetic to the GI tract there may be an increased incidence).

[0049] Adverse experiences (AE) information was assessed each morning at the time of dosing and again 24 hours following the last dose. Investigators and subjects were given diary cards detailing how to classify severity of AEs in order to standardise as much as possible across all 6 centres.

[0050] Of the 485 evaluable subjects, 18 (3.7%) withdrew, 17 because of adverse events. Subjects with nausea/ vomiting on the day of withdrawal were more common on (b) than either of (c) and (d). [0051] The incidence of nausea/vomiting and diarnhoea is shown in the table below.

	(b)	(c)	(d)	Placebo
Incidence of	59%	49%	39%	13%
nausea				
Incidence of	15%	21%	20%	7%
diarrhoea		l	l	1

[0652] The incidence of nausea was increased for both (b) and placebo compared to the expected rates of approximately 25% and 5% respectively for volunteers at these dosages for 5 days duration. The overall incidence of nausea was less on (c) and (c) than on (b). The severity of nausea was also decreased as shown in the next table.

Nausea severity	(b)	(c)	(d)	Placebo
None	50 (41%)	63 (52%)	74 (61 %)	104 (87%)
Mild	45 (37%)	40 (33%)	30 (25%)	16 (13%)
Moderate	21 (17%)	17 (14%)	15 (12%)	0 (0%)
Severe	6 (5%)	1 (1%)	3 (2%)	0 (0%)

[0053] Severity of diarrhoea is reported in the table below:

Severity of diarrhoea	(b)	(c)	(d)	Placebo
None	104 (85%)	95 (79%)	97 (80%)	112 (93%)
Mild	16 (13%)	16 (13%)	16 (13%)	8 (7%)
Moderate	1 (1%)	8 (7%)	9 (7%)	0 (0%)
Severe	1 (1%)	2 (2%)	0 (0%)	0 (0%)

[0054] In conclusion, there appears to be a trend for (c) to reduce the incidence of nauses and the dropout rate due to adverse events in comparison to (b), but analysis of the results was complicated by a statistically significant to transfer by-centre difference, (d) shows a harbing in the dropout rate and a fall in incidence of nauses of 20% (a proportional fall of 33%). In addition there is a reduction in severity of nauses of those build/builds who report nauses on (c) and (d). There is an increase in incidence of diarntoes on both of (c) and (d) in relation to (b), but this is confined to an increase in the number of individuals reporting moderate diarntoea and there is no increase in those with severe diarntoea.

Claims

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- A controlled release and delayed release formulation which is adapted or intended for oral administration and which contains paroxetine hydrochloride.
 - A controlled release and delayed release formulation according to claim 1 wherein paroxetine hydrochloride is in the form of paroxetine hydrochloride hemihydrate.
- 3. A tablet formulation which is a system for the controlled and delayed release of paroxeline hydrochloride, having (a) a desposit-core comprising 28.61 mg paroxeline hydrochloride, 18.75 mg methocal KMI, 79.14 mg lactose monohydrate, 2.50 mg polyvinylyprotidore, 1.25 mg magnesium stearate, and 0.50 mg syloid 244; and (b) a support-jainform applied to said deposit-core comprising 15.4 mg Comprisil 88.30, 30.0 mg lactose monothydrate, 4.00 mg polyvinylyprotidore, 9.80 mg magnesium stearate, 29.32 mg methocel ES, 0.32 mg Syloid 244, and 0.02 mg in on oxide and (c) an entire coatific comprising 13.27 mg Euradras 3.31 mg Lactose Lactose 1.32 mg methocal ES, 0.32 mg Syloid 244, and 0.02 mg in oxide and (c) an entire coatific comprising 13.27 mg Euradras 3.31 mg Lactose 1.32 mg method 13.32 mg and 13.32 mg Lactose 1.33 m



FUROPEAN SEARCH REPORT

Application Number EP 03 07 8259

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ANNEX TO THE EUROPEAN SEARCH REPORT ON EUROPEAN PATENT APPLICATION NO.

EP 03 07 8259

This ennex lists the patent family members relating to the patent documents cited in the above-mentioned European search report. The members are as contained in the European Petent Office EUP file on The European Petent Office is in new jack for three particulars which are merely given for the purpose of information.

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